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MANUAL MERCK

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DIAGNOSIS AND THERAPY

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FOREWORD

With this edition, The Merck Manual celebrates its 100th birthday. When the editors of the 1st Edition produced their 192-page compendium, they could not have realized the extent to which medical knowledge would explode over the next century. The Merck Manual now fills 2,655 pages and covers countless diseases that were not known 100 years ago. A brief review of medical practice as reflected in The Merck Manual during the past century follows on page vii.

Although the knowledge of medicine has grown, the goal of The Merck Manual has not changed—To provide useful clinical information to practicing physicians, medical students, interns, residents, nurses, pharmacists, and other health care professionals in a concise, complete, and accurate manner. The Merck Manual continues to cover all the subjects expected in a textbook of internal medicine as well as detailed information on pediatrics, psychiatry, obstetrics, gynecology, dermatology, pharmacology, ophthalmology, otolaryngology, and a number of special subjects. The Merck Manual quickly provides information that helps practitioners achieve optimal care. The more specialized the practice of medicine becomes, the more important such information becomes. Specialists as well specialises must at some time quickly access information about other specialises.

The 17th edition of The Merck Manual is the culmination of an arduous but rewarding 7-year enterprise. Every topic has been updated, and many have been completely rewritten. Topics new to this edition include hand disorders, prion diseases, death and dying, probabilities in clinical medicine, multiple chemical sensitivity, chronic fatigue syndrome, rehabilitation, smoking cessation, and drug therapy in the elderly, among others. The members of the Editorial Board, special consultants, and contributing authors are listed on the following pages with their affiliations. They deserve a degree of gratitude that cannot be adequately expressed here, but we know they will feel sufficiently rewarded if their efforts serve your needs.

Because of the extensive subject matter covered and a successful tradition developed through trials of successes and failures, *The Manual Manual* has some unique characteristics. We urge readers to spend a few minutes reviewing the Guide for Readers (p. xii), the Table of Contents at the beginning of each section (indicated by a thumb tab), and the Index (p. 2657). Subject headings within each section, internal headings within a subject discussion, and boldfaced terms in the text form an outline intended to help with use of the text.

We hope this edition of *The Merck Manual* will serve as an aid to you, our readers, compatible with your needs and worthy of frequent use. Suggestions for improvements will be warmly welcomed and carefully considered.

MARK H. BEERS, M.D., and ROBERT BERKOW, M.D., Editors

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DRUG INPUT AND DISPOSITION

Drugs are almost always compounds foreign to the body. As such, they, unlike endogenous substances, are not continually being formed and eliminated. Drug absorption, bioavallability, distribution, and elimination are therefore determinants of onset, duration, and intensity of drug effect.

ABSORPTION

mulated to be administered by various Drug products—the actual dosage forms (eg, tablets, capsules, solutions), consisting of the drug plus other ingredients—are for-A prerequisite to absorption is drug dissolution. Solid drug products (eg, tablets) disintegrate and deaggregate, but absorption can Drug absorption is determined by physicochemical properties of drugs, their foristration site to the systemic circulation. mulations, and routes of administration. routes, including oral, buccal, sublingual, rectal, parenteral, topical, and inhalational. Process of drug movement from the admin occur only after drugs enter solution.

Fransport Across Cell Membranes

a drug must traverse several semipermeable cell membranes before reaching the systemic circulation. These membranes are biologic barriers that selectively inhibit the posed primarily of a bimolecular lipid matrix, containing mostly cholesterol and phospholipids. The lipids provide stability to the membrane and determine its permeability characteristics. Globular proteins of various sizes and composition are embedded in the matrix; they are involved in transport and function as receptors for cellular regulation. Drugs may cross a biologic barrier by passive diffusion, facilitated passive diffusion, active passage of drug molecules and are com-When given by most routes (excluding IV), transport, or pinocytosis.

systemic circulation and distributed into a membrane by simple diffusion from a region of high concentration (eg, GI fluids) to one drug molecules are rapidly removed by the Passive diffusion: In this process, transport across a cell membrane depends on the concentration gradient of the solute. Most drug molecules are transported across a of low concentration (eg, blood). Because

producing a large gradient. The diffusion the area of the absorptive surface. Because large volume of body fluids and tissues, drug rate is directly proportional to the gradient but also depends on the molecule's lipid solubility, degree of ionization, and size and on the cell membrane is lipoid, lipid-soluble drugs diffuse more rapidly than relatively lipid-insoluble drugs. Small molecules tend to penetrate membranes more rapidly than concentration in blood is initially low compared with that at the administration site large ones.

that in the stomach. For a weak base with a pK_n of 4.4, the outcome is reversed. Thus theoretically, weakly acidic drugs (eg, aspiquinidine). However, whether a drug is concentration of ionized drug in the plasma would then be about 1000 times greater than rin) are more readily absorbed from an acid medium (stomach) than are weak bases (eg, on the cell membrane surface. Thus, drug penetration may be attributed mostly to the drug across a membrane at equilibrium is determined by the drug's pK_a (the pH at ized forms of the drug are equal) and the pH gradient, when present. Fór a weak acid, the higher the pH, the lower the ratio of un-ionthe ratio of un-ionized to ionized forms for a weak acid (eg, with a pK_a of 4.4) is 1:1000; in gastric fluid (pH, 1.4), the ratio is reversed (1000:1). When the weak acid is given orally, the concentration gradient for un-ionized drug between stomach and plasma tends to be large, favoring diffusion through the gastric mucosa. At equilibrium, the concentrations of un-ionized drug in the stomach and in the plasma are equal because only un-ionized drug can penetrate the membranes; the Most drugs are weak organic acids or bases, existing in un-ionized and ionized ionized form cannot penetrate the cell ubility and high electrical resistance, resulting from its charge and the charged groups un-ionized form. Distribution of an ionizable which concentrations of un-ionized and ionized to ionized forms. In plasma (pH, 7.4), forms in an aqueous environment. The unfuses readily across cell membranes. The membrane easily because of its low lipid solionized form is usually lipid soluble and dif

in the small intestine (see Oral Administra-

energy expenditure, and transport against a The carrier transports only substrates with a theory is that a carrier component combines reversibly with the substrate molecule at the strate complex diffuses rapidly across the membrane, releasing the substrate at the interior surface. Carrier-mediated diffusion is relatively specific molecular configuration, and the process is limited by the availability of carriers. The process does not require tain molecules (eg, glucose), the rate of pected from their low lipid solubility. One cell membrane exterior, and the carrier-subcharacterized by selectivity and saturability: membrane penetration is greater than ex-Facilitated passive diffusion: For cerconcentration gradient does not occur.

against a concentration gradient. Active transport appears to be limited to drugs Substrates may accumulate intracellularly stances. These drugs are usually absorbed from sites in the small intestine. Active transport processes have been identified for varstructurally similar to endogenous subacterized by selectivity and saturability and Active transport: This process is charrequires energy expenditure by the cell.

taches and moves to the cell interior. This hure. Pinocytosis probably plays a minor role nates, encloses the fluid or particles, then mechanism also requires energy expendiious ions, vitamins, sugars, and amino acids.

Pinocytosis: Fluid or particles are enguled by a cell. The cell membrane invagifuses again, forming a vesicle that later dein drug transport, except for protein drugs.

Oral Administration

tion is explained by the larger surface area and greater permeability of the membranes branes. However, the apparent contradicun-ionized drug more readily crosses memadministration is confounded by differences in luminal pH along the GI tract, surface area per luminal volume, blood perfusion, the presence of bile and mucus, and the nature of epithelial membranes. Acids are absorbed faster in the intestine than in the stomach, apparently contradicting the hypothesis that route, absorption refers to the transport of drugs across membranes of the epithelial cells in the GI tract. Absorption after oral For oral administration, the most common in the small intestine.

contact is usually too brief, even for drugs cur. A drug placed between the gums and tongue (sublingual administration) is re-tained longer so that absorption is more The oral mucosa has a thin epithelium and a rich vascularity that favors absorption, but cheek (buccal administration) or under the in solution, for appreciable absorption to occomplete.

(eg, penicillin G), or have little or no effect Drugs that affect gastric emptying (eg, parasympatholytic drugs) affect the absorption of drug absorption), explaining why some when a rapid onset of action is desired. Food may enhance the extent of absorption for poorly soluble drugs (eg, griseofulvin), reduce it for drugs degraded in the stomach is limited. Absorption of virtually all drugs is faster from the small intestine than from the stomach. Therefore, gastric emptying is the rate-limiting step. Food, especially fatty foods, slows gastric emptying (and the rate drugs should be taken on an empty stomach there is usually relatively short, absorption The stomach has a relatively large epithecous layer and the time that the drug remains lial surface, but because it has a thick murate of other drugs.

intraluminal pH is 4 to 5 in the duodenum but becomes progressively more alkaline, flora may inactivate certain drugs, reducing dient across the intestinal mucosa and decrease absorption by passive diffusion. (Decreased peripheral blood flow also alters area for drug absorption in the GI tract. The approaching 8 in the lower ileum. GI microtheir absorption. Decreased blood flow (eg, in shock) may lower the concentration gra-The small intestine has the largest surface drug distribution and metabolism.)

biotics). For such drugs, transit may be too or that are too polar (ie, poorly lipid-soluble) to cross membranes readily (eg, many anti-Intestinal transit time can influence drug absorption, particularly for drugs that are absorbed by active transport (eg, B vitamins), that dissolve slowly (eg, griseofulvin), rapid for absorption to be complete.

tinues for > 6 h, the time for transit to the sorption may occur primarily in the large intestine, particularly when drug release con-For controlled-release dosage forms, ablarge intestine.

Almorption from solution: A drug given orally in solution is subjected to numerous (H secretions and, to be absorbed, must sur-

acidic or basic, most of its absorption occurs

is stable in the enteral environment, little of it remains to pass into the large intestine. Drugs with low lipophilicity (ie, low memsides, are absorbed slowly from solution in drugs, absorption in the large intestine is exbrane permeability), such as aminoglycothe stomach and small intestine; for such pected to be even slower because the surface area is smaller. Consequently, these drugs are not candidates for controlled release.

Absorption from solid forms: Most drugs are given orally as tablets or capsules primarily for convenience, economy, stability, and patient acceptance. These products must disintegrate and dissolve before abcreases the drug's surface area in contact with GI fluids, thereby promoting drug dissolution and absorption. Disintegrants and sorption can occur. Disintegration greatly insurfactants, binders, dispersants) are often added during manufacture to facilitate these tion rate by increasing the wetability, solubility, and dispersibility of the drug. Disinother excipients (eg. diluents, lubricants, processes. Surfactants increase the dissolutegration of solid forms may be retarded by plied to protect the tablet from the digestive processes of the gut. Hydrophobic lubricants (eg, magnesium stearate) may bind to the excessive pressure applied during the tableting procedure or by special coatings apactive drug and reduce its bioavailability.

Dissolution rate determines the availability of the drug for absorption. When slower than absorption, dissolution betion can be controlled by manipulating the tion rate is affected by whether the drug is in salt, crystal, or hydrate form. The Na salts formulation. For example, reducing the particle size increases the drug's surface area, dissolve faster than their corresponding free comes the rate-limiting step. Overall absorpthus increasing the rate and extent of GI absorption of a drug whose absorption is normally limited by slow dissolution. Dissoluof weak acids (eg, barbiturates, salicylates) acids regardless of the pH of the medium. Certain drugs are polymorphic, existing in amorphous or various crystalline forms. sorbed to be clinically useful. A hydrate is formed when one or more water molecules Chloramphenicol palmitate has two forms, but only one sufficiently dissolves and is abcombine with a drug molecule in crystal

form. The solubility of such a solvate may markedly differ from the nonsolvated form; eg, anhydrous ampicillin has a greater rate of dissolution and absorption than its corresponding trihydrate.

Parenteral Administration

stream (usually IV) ensures delivery of the delivery of the entire dose is not ensured if a route requires movement through one or more biologic membranes to reach the systemic circulation (IM or sc injection). For membranes is so slow that after IM or sc administration, most absorption occurs via Direct placement of a drug into the blood dose to the systemic circulation. However, protein drugs with a molecular mass > 20,000 g/mol, movement across capillary the lymphatic system by default. In such cases, the delivery rate to systemic circulation is slow and often incomplete because of first-pass metabolism by proteolytic enzymes in the lymphatics.

Because capillaries tend to be highly porous, perfusion (blood flow/gram of tissue) molecules. Thus, the injection site can markinto a site with poor blood flow can be much greatly affects the absorption rate of small edly influence a drug's absorption rate; eg, the absorption rate of diazepam injected IM slower than that after oral administration.

Absorption may be delayed or erratic when salts of poorly soluble acids and bases toin is a 40% propylene glycol solution of is absorbed, and the tissue fluids, acting as a are injected IM. The parenteral form of phenthe Na salt with a pH of about 12. When the rium between the ionized and free acid forms solution is injected IM, the propylene glycol buffer, decrease the pH, shifting the equilibof the drug. The poorly soluble free acid then precipitates. As a result, dissolution and absorption take 1 to 2 wk to occur.

Controlled-Release Forms

Controlled-release dosage forms are designed to reduce dosing frequency and to reduce fluctuation in plasma drug concentration, providing a more uniform the rapeutic effect. Less frequent dosing is more convenient and may improve patient compliance. These dosage forms are suitable for drugs that otherwise require frequent dosing because elimination half-life and duration of effect are short.

Oral controlled-release forms are often designed to maintain therapeutic drug concentrations for ≥ 12 h. The absorption rate can wax or other water-insoluble material, by the GI tract, or by complexing the drug with be controlled by coating drug particles with embedding the drug in a matrix from which it is released slowly during transit through ion-exchange resins.

Transdermal controlled-release forms are designed to provide drug release for expregnated polymer bonded to an adhesive tended periods; eg, clonidine diffusion through a membrane provides controlled drug delivery for 1 wk, and nitroglycerin-imfor 24 h. Drugs for transdermal delivery must have suitable skin penetration characterisbandage provides controlled drug delivery tics and high potency because the penetration rate and area of application are limited.

els. For antimicrobials, relatively insoluble extended periods. For others, suspensions sulin injected in crystalline suspensions) are formulated. Amorphous insulin, with a high Many nonintravenous parenteral preparations are formulated to sustain blood levsalts (eg, penicillin G benzathine) injected IM provide therapeutic concentrations for or solutions in nonaqueous vehicles (eg, insurface area for dissolution, has a rapid onset and short duration of action.

BIOAVAIL A BILITY

Extent to which—and sometimes rate at which-the active moiety (drug or methereby gaining access to the site of actabolite) enters systemic circulation,

pend on its design and manufacture) can largely determine drug bioavailability. Dificance. Thus, the concept of equivalence clinical decisions. Chemical equivalence The physicochemical properties of a drug erties of the dosage form (which partly deferences in bioavailability among formulaamong drug products is important in making refers to drug products that contain the same compound in the same amount and that meet current official standards; however, inactive Bioequivalence refers to chemical equivations of a given drug can have clinical signifingredients in drug products may differ. govern its absorptive potential, but the prop

fers to drug products that, when administered to the same person in the same dosage lents that, when administered to the same person in the same dosage regimen, result in regimen, provide essentially the same therapeutic effect or toxicity. Bioequivalent products are expected to be therapeutically equivalent concentrations of drug in blood and tissues. Therapeutic equivalence reequivalent,

is stabilized on one formulation is given a Therapeutic problems (eg, toxicity, lack of during long-term therapy when a patient who efficacy) are encountered most frequently nonequivalent substitute (as for digoxin or phenytoin).

Sometimes therapeutic equivalence may ability. For example, the therapeutic index (ratio of the maximum tolerated dose to the minimum effective dose) of penicillin is so penicillin products may not affect therapeutic efficacy or safety. In contrast, bioavailability differences are important for a drug wide that moderate blood concentration differences due to bioavailability differences in be achieved despite differences in bioavailwith a relatively narrow therapeutic index.

The physiologic characteristics and comorbidities of the patient also affect bioavailability.

when a drug is absorbed completely, it may be absorbed too slowly to produce a therapeutic blood level quickly enough or so rap-Absorption rate is important because even idly that toxicity results from high drug concentrations after each dose.

Causes of Low Bioavailability

tered drugs is not always complete. Before crosses membranes, absorption tends to be complete, but absorption of orally adminisreaching the vena cava, a drug must move When a drug rapidly dissolves and readily down the GI tract and pass through the gut wall and liver, common sites of drug metabolism (see Ch. 43); thus, a drug may be metabolized (first-pass metabolism) before it For such drugs (eg, isoproterenol, norepinephrine, testosterone), extraction in these lissues is so extensive that bioavailability can be measured in the systemic circulation. Many drugs have low oral bioavailability because of extensive first-pass metabolism. is virtually zero. For drugs with an active metabolite, the therapeutic consequence of Itrst-pass metabolism depends on the contri-



是是我们的,但是我们的人,我们可以一个时间的一个人的,我们也是我们的一个人的,我们也是我们的一个人,我们也是我们的人的,我们也是我们的人,我们们是什么一个人,

butions of the drug and the metabolite to the desired and undesired effects.

Low bioavailability is most common with fect bioavailability when absorption is slow slowly absorbed drugs. More factors can aforal dosage forms of poorly water-soluble, or incomplete than when it is rapid and complete, so slow or incomplete absorption often leads to variable therapeutic responses.

Insufficient time in the GI tract is a common cause of low bioavailability. Ingested drug is exposed to the entire G1 tract for no more than 1 to 2 days and to the small intestine for only 2 to 4 h. If the drug does not dissolve readily or cannot penetrate the epithelial membrane (eg, if it is highly ionized and polar), time at the absorption site may be insufficient. In such cases, bioavailability Age, sex, activity, genetic phenotype, stress, disease (eg, achlorhydria, malabsorption tends to be highly variable as well as low. syndromes), or previous GI surgery can affect drug bioavailability.

Reactions that compete with absorption acid or digestive enzymes (eg, penicillin and plex formation (eg, between tetracycline and polyvalent metal ions), hydrolysis by gastric chloramphenicol palmitate hydrolysis), conjugation in the gut wall (eg, sulfoconjugation of isoproterenol), adsorption to other drugs can reduce bioavailability. They include com-(eg, digoxin and cholestyramine), and metabolism by luminal microflora.

Assessment of Bloavailability

Assessment of bioavailability from plasma termining the maximum (peak) plasma drug with the extent of absorption; the peak is reached when the drug elimination rate tration can be misleading, because drug elimination begins as soon as the drug enters concentration—time data usually involves deplasma drug concentration occurs (peak time), and the area under the plasma con- The plasma drug concentration increases minations based on the peak plasma concenconcentration, the time at which maximum equals absorption rate. Bioavailability deterthe bloodstream. The most widely used genthe slower the absorption, the later the peak time. However, peak time is often not a good statistical measure because it is a discrete centration-time curve (AUC-see Fig. 298value that depends on frequency of blood eral index of absorption rate is peak time;



FIG. 298-1. Representative plasma concentration—time relationship after a single oral dose of a hypothetical drug. Area under the plasma concentration-time curve is indicated by shading.

concentrations near the peak, on assay reproducibility.

total amount of unchanged drug that reaches AUC is the most reliable measure of bioavallability. It is directly proportional to the the systemic circulation. For an accurate measurement, blood must be sampled frequently over a long enough time to observe virtually complete drug elimination. Drug products may be considered bioequivalent in extent and rate of absorption if their plasma-level curves are essentially superimposable. Drug products that have similar AUCs but differently shaped plasma-level curves are equivalent in extent but differ in their absorption rate-time profiles.

ity may be assessed after single or repetitive Single vs. multiple doses: Bioavailabil-(multiple) dosing. More information about rate of absorption is available after'a single dose than after multiple dosing. However, multiple dosing more closely represents the trations are usually higher than those after a single dose, facilitating data analysis. After usual clinical situation, and plasma concenmultiple dosing at a fixed-dosing interval for cour or five elimination half-lives, the blood drug concentration should be at steady state the amount absorbed equals the amount eliminated within each dosing interval). The extent of absorption can then be analyzed by measuring the AUC during a dosing interval. Measuring the AUC over 24 h is probably preferable because of circadian variations in physiologic functions and because of possible variations in dosing intervals and absorption rates during a day.

For drugs excreted primarily unchanged in urine, bioavailability can be estimated by

sampling and, in the case of relatively flat

unchanged drug recovered from unine over measuring the total amount of drug excreted after a single dose. Ideally, urine is collected over a period of 7 to 10 elimination half-lives for complete urinary recovery of the absorbed drug. Bioavailability may also be assessed after multiple dosing by measuring 24 h under steady-state conditions.

DISTRIBUTION

After a drug enters the systemic circula-Distribution is generally uneven because of differences in blood perfusion, tissue binding, regional pH, and permeability of cell tion, it is distributed to the body's tissues. membranes.

istics between blood and tissue. Distribution equilibrium (when entry and exit rates are same) between blood and tissue is reached more rapidly in richly vascularized areas than in poorly perfused areas, unless diffusion across membrane barriers is the 늃 tained, drug concentrations (bound and uncretion occur simultaneously with distribu-The entry rate of a drug into a tissue depends on the rate of blood flow to the tissue, on tissue mass, and on partition characterplasma concentration. Metabolism and extion, making the process dynamic and comrate-limiting step. After equilibrium is bound-see below) in tissues and extracellular fluids are reflected by plex (see also Ch. 299)

Apparent Volume of Distribution

volume required to contain the drug in the body at the same concentration as in plasma). This parameter provides a refer-The volume of fluid into which a drug appears to be distributed or diluted is called the apparent volume of distribution (the fluid ence for the plasma concentration expected for a given dose and for the dose required to produce a given concentration. However, it provides little information about the specific pattern of distribution. Each drug is uniquely distributed in the body. Some drugs go into fat, others remain in the ECF, and still others are bound avidly to specific tissues, commonly liver or kidney.

Many acidic drugs (eg, warfarin, salicylic acid) are highly protein-bound and thus have small apparent volume of distribution. Many basic drugs (eg, amphetamine, meper-

idine) are avidly taken up by tissues and thus have an apparent volume of distribution CHAPTER 298 - DRUG INPUT AND DISPOSITION / 2561 larger than the volume of the entire body.

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Binding

The extent of drug distribution into tissues depends on the extent of plasma protein and tissue binding.

Albumin, \$\alpha_1\$-acid glycoprotein, and lipoproteins are most important. Acidic drugs are Plasma protein binding: Drugs are transported in the bloodstream partly in solution as free (unbound) drug and partly bound to reversible interaction between a drug and the plasma protein to which it binds, as governed by the law of mass action. Many blood components (eg, plasma proteins, blood cells). The ratio of bound to unbound drug in plasma is mainly determined by the plasma proteins can interact with drugs. generally bound more extensively to albumin, and basic drugs to α_l -acid glycoprotein and/or lipoproteins (see TABLE 298-1)

bound. Plasma protein binding influences Only unbound drug is thought to be availcur. Therefore, the unbound drug concentration may be more closely related to drug concentration at the active site and to drug effects, often making the fraction unbound able for passive diffusion to extravascular or tissue sites where pharmacologic effects oc-(ratio of unbound to total concentrations) a more useful parameter than the fraction distribution and the apparent relationship between pharmacologic activity and total

TABLE 298-1. EXTENT OF BINDING IN PLASMA FOR SELECTED DRUGS

ין אוונטין י	LASIMA FOR SELECTED DRUGS	D DREES
Drug	% Bound	% Unbound
Warfarin	99.5	0.5
Diazepam	66	_
Furosemide	96	4
Dicloxacillin	94	9
Propranolol*	83	7
Phenytoin	68	11
Quinidine*	71	53
Lidocaine*	51	49
Digoxin	22	75
Gentamich	က	26
Atenolol	0~	~100

^{*}Significant hinding to a1-acid glycoprotein and/or lipoproteins.



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plasma drug concentration. At high drug concentrations, the amount of bound drug approaches an upper limit depending on the number of available binding sites, resulting displacement interactions among drugs (see in saturability. Saturability is the basis of DRUG INTERACTIONS in Ch. 301).

Tissue binding: Drugs bind to many substances other than proteins. Binding may be very specific, as when chloroquine binds with nucleic acids. Binding usually occurs when a drug associates with a macromolecule in an aqueous environment but may occur when a drug is partitioned into body fat. Because fat is poorly perfused, equilibration time is long, especially if the drug has a high affluity for fat.

the sojourn of drug in plasma and drug action because the tissues release stored drug as in tissues or body compartments can prolong Drug reservoir: Accumulation of drugs the plasma concentration declines. Location of the active site and relative differences in tissue distribution can also be important. For the anesthetic thiopental, storage in tissue reservoirs initially shortens the drug effect but after repeated administration prolongs it. Thiopental is highly lipid soluble and rapidly distributes to the brain after a single IV injection. After a single dose, thiopental concentration in the brain increases for a few minutes, then declines parallel with the plasma concentration. Anesthesia ends rapidly as the drug redistributes to more slowly centration is monitored long enough, a third phase of distribution, in which the drug is perfused tissues. However, if plasma conguished. With continued administration of slowly released from fat, can be distinfat, resulting in prolongation of anesthetic thiopental, large amounts may be stored in plasma concentrations.

Some drugs accumulate, producing higher concentrations in cells than in ECF, most phospholipids, or nucleic acids. Antimalarial tions within WBCs and liver cells thousands stored drug is in equilibrium with drug in plasma and moves into plasma as the drug is commonly because they bind with protein drugs (eg, chloroquine) produce concentraof times higher than those in plasma. The eliminated from the body.

lood-Brain Barrier

Drugs reach the CNS via brain capillaries and via CSF, Although the brain receives

about 1/6 of cardiac output, distribution of drugs to brain tissue is restricted. Some lipidsoluble drugs (eg, thiopental) enter the brain and exert their pharmacologic effects rapidly, but many drugs, particularly the more which appear to be more tightly joined to one another than are those of other capillarwater-soluble drugs, enter the brain slowly. The endothelial cells of the brain capillaries, ies, contribute to the slow diffusion of watersoluble drugs. Another barrier to watersoluble drugs is the glial connective tissue cells (astrocytes), which form an astrocytic sheath close to the basement membrane of the capillary endothelium. The capillary endothelium and the astrocytic sheath form the blood-brain barrier. Because the capillary wall rather than the parenchymal cell forms the barrier, the brain's permeability charac-Thus, polar compounds cannot enter the teristics differ from those of other tissues. brain but can enter the interstitial fluids of most other tissues. The observation that polar dyes enter most tissues but not the CNS led to the concept of the blood-brain barrier.

via the choroid plexus, entering brain tissue Drugs may enter ventricular CSF directly by passive diffusion from CSF. Also in the choroid plexus, organic acids (eg. penicillin) are actively transported from CSF to blood.

The drug penetration rate into the CSF or into other tissue cells is determined mainly of ionization, and the lipid-water partition by the extent of protein binding, the degree coefficient of the drug. The penetration rate into the brain is slow for highly proteinbound drugs and can be so slow for the ionized form of weak acids and bases as to be virtually nonexistent.

meability is generally the major determinant of the drug distribution rate. However, for Because the CNS is so well perfused, perthe interstitial fluids of most tissues, perfusion is a major determinant. For poorly perfused tissues (eg, muscle, fat), distribution is very slow, especially if the tissue has a high affinity for the drug.

ELIMINATION

Sum of the processes of drug loss (metabolism and excretion) from the body.

METABOLISM

The liver is the principal site of drug metabolism (chemical alteration) in the body.

CHAPTER 298 - DRUG INPUT AND DISPOSITION / 2563

SELECTED DRUGS WITH THERAPEUTICALLY IMPORTANT METABOLITES TABLE 298-2.

Drug	Metabolite
Acetohexamide	Hydroxyhexamide
Amitriptyline	Nortriptyline
Aspirin*	Salicylic acid
Chloral hydrate*	Trichloroethanol
Chlordiazepoxide	Desmethylchlordiaze-
	poxide
Codeine	Morphine
Diazepam	Desmethyldiazepam
Flurazepam	Desethylflurazepam
Glutethimide	4-Hydroxyglutethimide
Imipramine	Desipramine
Lidocaine	Desethyllidocaine
Meperidine	Normeperidine
Phenacetin*	Acetaminophen
Phenylbutazone	Oxyphenbutazone
Prednisone*	Prednisolone
Primidone*	Phenobarbital
Procainamide	N-acetylprocainamide
Propranolol	4-Hydroxypropranolol

*Pro-drugs; metabolites are primarily responsible for their therapeutic effects. Some metabolites are pharmacologically stance that has an active metabolite is called a pro-drug, especially if designed to deliver active (see TABLE 298-2). An inactive subthe active moiety more effectively.

Pathways of Metabolism

thetic reactions. Phase II reactions involve Drug metabolism involves a wide range of chemical reactions, including oxidation, reduction, hydrolysis, hydration, conjugation, condensation, and isomerization. The enzymes involved are present in many tissues but generally are more concentrated in the liver. For many drugs, metabolism occurs in volve the formation of a new or modified functional group or a cleavage (oxidation, reduction, hydrolysis); these are nonsynconjugation with an endogenous compound (eg, glucuronic acid, sulfate, glycine) and are therefore synthetic reactions. Metabolites and more readily excreted by the kidneys (in urine) and the liver (in bile) thun those formed in nonsynthetic reactions, Some drugs undergo either phase I or pluse II retwo apparent phases. Phase I reactions informed in synthetic reactions are more polar

actions; thus, phase numbers reflect functional rather than sequential classification.

ADPH-cytochrome P-450 reductase, a Cytochrome P-450: The most important ochrome P-450, a microsomal superfamily if isoenzymes that transfer electrons and nereby catalyze the oxidation of many The electrons are supplied by avoprotein that transfers electrons from ADPH (the reduced form of nicotinamidedenine dinucleotide phosphate) to cytohrome P-450. Cytochrome P-450 enzymes unily, a letter for subfamily, and another ies are most important in mammalian 301). Genetic differences among patients nzyme system of phase I metabolism is cyre grouped into 14 mammalian gene famies that share sequence identity and 17 sub-amilies. They are designated by a root symol CYP, followed by an Arabic number for rabic number for the specific gene. Enymes in the 1A, 2B, 2C, 2D, and 3Ă subfamcytochrome P-450 enzymes are listed in TAetabolism; CYP1A2, CYP2C9, CYP2C19, YP2D6, and CYP3A4 are important in huian metabolism. The specificity of the enrmes helps explain many drug interactions. xamples of drugs that interact with specific BLE 298-3 (see also DRUG INTERACTIONS in Ch. may alter response.

Conjugation: Glucuronidation, the most that occurs in the liver microsomal enzyme glutamine or glycine produces conjugates common phase II reaction, is the only one system. Glucuronides are secreted in bile lized this way. Amino acid conjugation with meprobamate, and morphine are metaboeg, salicyluric acid formed from salicylic acid and glycine) that are readily excreted in urine but are not extensively secreted in bile. Acetylation is the primary metabolic pathphenolic or alcoholic groups and inorganic ily excreted in urine. Drugs that form sulfate way for sulfonamides. Hydralazine, isoniaand eliminated in urine. Chloramphenicol zid, and procainamide are also acetylated Sulfoconjugation is the reaction between sulfate, which is partially derived from sulfur-containing amino acids (eg, cysteine) The sulfate esters formed are polar and readconjugates include acetaminophen, estradiol, methyldopa, minoxidil, and thyroxine. Methylation is a major metabolic pathway Nincinamide and thiouracil are also methylfor inactivation of some catecholamines <u>=</u>

TABLE 298-3. SOME SUBSTANCES THAT INTERACT WITH CYTOCHROME P-450

		l beef	•	· · · · · · · · · · · · · · · · · · ·		
	Inducare	Charcoal-broiled beef	Rifampin	Rifampin	None known	Carbamazepine Phenobarbital
ENZYMES	Inhibitors	Furafylline	Sulfaphenazole Sulfinpyrazone	Tranylcypromine	Fluoxetine Quinidine	Ketoconazole Troleandomycin
	Substrates	Acetaminophen Estradiol Theophylline Verapanil	Dictofenac Phenytoin Piroxican Tetrahydrocannabinol Tolbutamide	Diazepam Hexobarbital Omeprazole Pentamidine Propranolol	Debrisoquin Desipramine Encainide Mexiletine Nortriptyline	Amiodarone Lovastatin Nifedipine Tamoxifen Terfenadine
	Enzyme	CYP1A2	CYP2C9	CYP2C19	CYP2D6	CYP3A4

Age-Related Changes

Because newborns have partially developed liver microsomal enzyme systems, they have difficulty metabolizing many drugs (eg. hexobarbital, phenacetin, amphetamine, chlorpromazine). In newborns, slower conversion to glucuronide can have serious effects. For example, equivalent mg/kg doses of chloramphenicol that are well tolerated by older patients can result in the gray baby syndrome and in prolonged elevated blood levels of chloramphenicol.

Elderly patients often have a reduced ability to metabolize drugs. The reduction varies depending on the drug and is not as severed that in newborns (see Ch. 304).

ndividual Variation

Betause of individual variation (see also VARIABILITY IN PARAMETER VALUES in Ch. 299), predicting the clinical response to a given dose of a drug is difficult. Some patients me-

abolize a drug so rapidly that therapeutically effective blood and tissue concentrations are not achieved; in others, metabolism may be so slow that usual doses produce toxic effects. For example, plasma phenytoin concentrations at steady state vary from $2.5 \text{ to} > 40 \text{ mg/L} (10 \text{ to} > 160 \text{ } \mu\text{mol/L}) \text{ in}$ different patients given a daily dose of 300 mg. Some variation is due to differences in available in the liver and to differences in the the amount of the key enzyme, CYP2C9, affluity of the enzyme for the drug. Genetic factors play a major role in determining ticularly chronic liver disease) and drug these differences. Concurrent diseases (parduction or inhibition of metabolism) also interactions (especially those involving incontribute.

Capacity Limitation

For almost any drug, the rate of metabolism of any enzyme in any given pathway

reaches an upper limit (capacity limitation). At therapeutic concentrations, usually only a small fraction of the enzyme sites are occupied, and the rate of metabolism increases with drug concentration. Occasionally, when most of the enzyme sites are occupied, the rate of nuctabolism does not increase in proportion to drug concentration. The result is capacity-limited metabolism. Phenytoin and alcohol have this type of metabolism, which helps explain the interpatient variability in phenytoin concentrations after a fixed duily dose of 300 mg.

EXCRETION

Process by which a drug or a metabolite is eliminated from the body without further chemical change.

The kidneys, which excrete water-soluble substances, are the major organs of excretion. The biliary system contributes to excretion to the degree that drug is not reabsorbed from the GI tract. Generally, the contribution of intestine, saliva, sweat, breast milk, and lungs to excretion is small, except for exhalution of volatile anesthetics. Although excretion via breast milk may not be important to the mother, it may be to the suckling infant (see Drucs in LACTATING MOTHERS in Ch. 256).

Renal Excretion

Glomerular filtration and tubular reabsorption: About 1/5 of the plasma reaching the glomerulus is filtered through pores in the glomerular endothelium; the remainder passes through the efferent arterioles surrounding the renal tubules. Drugs bound to plasma proteins are not filtered; only unbound drug is contained in the filtrate. The principles of transmembrane passage govern renal tubular reabsorption of drugs. Polar compounds and ions cannot diffuse back into the circulation and are excreted unless a specific transport mechanism for their reabsorption exists (eg, as for glucose, ascorbic acid, and B vitamins).

Effects of urine pff: The glomerular filtrate that enters the proximal tubule has the same pH as plasma, but the pH of voided urine varies from 4.5 to 8.0. This variation in pH may markedly affect the rate of drug excretion. Because un-ionized forms of non-polar weak acids and weak bases tend to be reabsorbed readily from tubular fluids, acid-

ification of urine increases reabsorption (ie, decreases excretion) of weak acids and decreases reabsorption (ie, increases excretion) of weak bases. The opposite occurs after alkalinization of urine.

In some cases of overdose, these principles may be applied to enhance the exerction of weak acids or bases. For example, alkalinization of urine increases the excretion of the weak acids phenobarbital and aspirin, and acidification may accelerate the excretion of bases, such as methamphetamine. The extent to which changes in urinary pll after the rate of drug elimination depends on the contribution of the renal route to total elimination as well as on the polarity of the un-ionized form and the degree of ionization of the molecule.

Tubular secretion: Mechanisms for active tubular secretion in the proximal tubule are important in the elimination of many drugs (eg, penicillin, mecanylamine, salicylic acid). This energy-dependent process may be blocked by metabolic inhibitors. When drug concentration is high, an upper limit for secretory transport can be reached; each substance has a characteristic maximum secretion rate (transport maximum).

Anions and cations are handled by separate transport mechanisms. Normally, the anion secretory system eliminates metabolites conjugated with glycine, sulfate, or glucuronic acid. Anionic compounds compete with one another for secretion. This competition can be used therapeutically; eg, probenecid blocks the normally rapid tubular secretion of penicillin, resulting in higher plasma penicillin concentrations for a longer time. Organic cations compete with each other but usually not with anions.

Age-related changes: With aging, renal drug excretion decreases (see Pharwacokiners in Ch. 304 and Table 304-1).

Billary Excretion

Drugs and their metabolites that are extensively excreted in bile are transported across the biliary epithelium against a concentration gradient, requiring active secretory transport. Secretory transport may approach an upper limit at high plasma concentrations of a drug (transport maximum), and substances with similar physicochemical properties may compete for excretion via the same mechanism.



Drugs with a mol wt $> 300 \,\mathrm{g/mol}$ (smaller molecules are generally excreted only in creted in bile. Conjugation, particularly with glucuronic acid, also leads to biliary excrenegligible amounts) and with both polar and lipophilic groups are more likely to be ex-

In the enterohepatic cycle, a drug secreted in bile is reabsorbed from the intes-

tine. Drug conjugates secreted into the tines

substances from the body only to the exami when they are hydrolyzed and the drug is tine also undergo enterohepatic براالله reabsorbed. Biliary excretion eliminants that enterohepatic cycling is incompleकां ह when some of the secreted drug is not math sorbed from the intestine.

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Study of the time course of a drug and its metabolites in th body after administration by any route.

An appropriate response to a drug requires the appropriate concentration of drug at the site of action. The dosage regimen required to attain and maintain the appropriate concentration depends on pharmacokinetics. The appropriate concentration and dosage regimen depend on the patient's clinical state, severity of the disorder, presence of concurrent disease, use of other drugs, and

ministration must be based on each patient's ing dosage until the therapeutic objective is Because of individual differences, drug adneeds-traditionally, by empirically adjustmet. This approach is frequently inadequate tively, a drug can be administered according because optimal response may be delayed or serious toxic reactions may occur. Alternato its expected absorption and disposition 298) in a patient, and dosage can be adjusted and drug effects. This approach requires distribution and elimination—see also Ch. by monitoring plasma drug concentration knowledge of the drug's pharmacokinetics as a function of the patient's age and weight and the kinetic consequences of concurrent diseases (eg, renal, hepatic, or cardiovascu. lar disease or a combination of diseases).

PHARMACOKINETIC PARAMETERS BASIC

The pharmacokinetic behavior of most drugs can be summarized by the following parameters, whose formulas are listed in IABLE 299-1. The parameters are constants,

although their values may differ from pathens to patient and in the same patient under dif ferent conditions.

Bioavailability expresses the extent of drug absorption into the systemic circulates maximum concentration occurs (peul see Ch. 298). The absorption rate constant expresses the speed of absorption (peak) concentration, the time at which the time), and the area under the concentraixm time curve (AUC) after a single oral done: During long-term drug therapy, the extens of absorption is the more important measure ment because average concentration de pends on it; the degree of fluctuation is re-These parameters influence the maximm lated to the absorption rate constant.

The apparent volume of distribution is the amount of fluid that would be required to contain the drug in the body at the sazar concentration as in the blood or plasma in can be used to estimate the dose required to produce a given concentration and the con centration expected for a given dose. The unbound concentration is closely associand useful measure, particularly when plasma protein binding is altered-eg, by hypoalboume of distribution and the unbound fraction with drug effects, so unbound fraction is a placement interactions. The apparent we in plasma are the most widely used param minemia, renal or hepatic disease, or daeters for drug distribution (see Ch. 298).

The rate of elimination of a drug from the body varies with the plasma concentration The parameter relating elimination rate to which equals renal clearance plus extrarenal plasma concentration is total clearance,

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1481E 299-1. FORMULAS DEFINING BASIC PHARMACOKINETIC PARAMETERS

		Malt 222-1. Formation behavior from the contraction of the contraction
Cathegory	Parameter	Formula
-фонду-	Absorption rate contant	 Rate of drug absorption + Amount of drug remaining to be absorbed
	Bioavailability	= Amount of drug absorbed + Drug dose
fluaribu-	Apparent volume of distribution	= Amount of drug in body + Plasma drug concentration
	Unbound fraction	 Plasma concentration of unbound drug + Plasma drug concentration
Elmina- tion	Rate of elimination	= Renal excretion + Extrarenal (usually metabolic) elimination
	Clearance	 Rate of drug elimination + Plasma drug concentration
	Renal clearance	 Rate of renal excretion of drug + Plasma drug concentration
	Metabolic clearance	 Rate of drug metabolism + Plasma drug con- centration
,	Fraction excreted unchanged	 Rate of renal excretion of drug + Rate of drug elimination
	Elimination rate constant	 Rate of drug elimination + Amount of drug in body
	-	= Clearance + Volume of distribution
	Biologic half-life	= 0.693 + Elimination rate constant

ametabolic) clearance (see also Estimation of Parameter Values in Ch. 303).

faction indicates that hepatic metabolism is The fraction excreted unchanged helps the likely mechanism of elimination and that pepatic disease may therefore affect drug dinination. Renal diseases produce greater effects on the kinetics of drugs with a high patic diseases on drug elimination. A low assess the potential effect of renal and he fraction excreted unchanged.

and a drug is given orally, part of the dose may be metabolized as it passes through the the liver or gut wall, oral bioavailability is low, sometimes precluding oral administrathan an equivalent parenteral dose. Drugs with extensive first-pass metabolism include uprenolol, hydralazine, isoproterenol, lido-The extraction rate of a drug from the slood by an eliminating organ, such as the to the organ. Thus, clearance has an upper limit, based on drug delivery and hence on f extraction (clearance) of a drug is high in tion or requiring an oral dose much larger her, cannot exceed the rate of drug delivery blood flow to the organ. Furthermore, when usues to the systemic circulation; this proces is called first-pass metabolism. Thus, the eliminating organ is the liver or gut wall

troglycerin, propranolol, testosterone, and caine, meperidine, morphine, nifedipine, ni-

verapamil.

tion of how a drug is cleared from the blood by the eliminating organs and how the drug The elimination rate constant is a func-

quired for the plasma drug concentration or the amount of drug in the body to decrease by 50%. For most drugs, half-life remains the same regardless of how much drug is in the distributes throughout the body.

Half-life (elimination) is the time rebody. Exceptions include phenytoin, theophylline, and heparin.

time a drug molecule remains in the body after rapid IV injection. Like clearance, its value is independent of dose. After an IV measure of drug elimination, is the average Mean residence time (MRT), another bolus,

$$MRT = \frac{AUMC}{ATIC}$$

the plasma concentration-time curve. For a drug with one-compartment distribution AUMC is the area under the first moment of characteristics, MRT equals the reciprocal of the elimination rate constant.





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DRUG ADMINISTRATION

doses are described below, using throphylcially children. In this example, the drug is given to a 70-kg patient (patient A) who has 1.0; absorption rate constant, 1.0/h; apparent ng a drug in a single dose (IV or oral), by constant-rate infusion, and in multiple oral line (given as aminophylline) as an example. The metabolism of theophylline is concentration-dependent in some persons, especoncentration-independent metabolism and volume of distribution, 0.5 L/kg; clearance, The kinetic consequences of administer the following parameters: bioavailability 43 mL/h/kg; and half-life, 8 h.

Single Dose

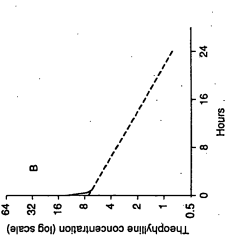
dose of aminophylline (hydrous form is 80% mated from the half-life; every 8 h, the con-Intravascular: After a single 320-mg IV 299-1), the predicted initial plasma concentration of theophylline is 7.3 mg/L (41 μ mol/L)—ie, dose (256 mg) divided by apparent volume of distribution (0.5 L/kg \times 70 kg = 35 L). The subsequent decline is estitheophylline) is given to patient A (see F10. centration decreases by a factor of 2.

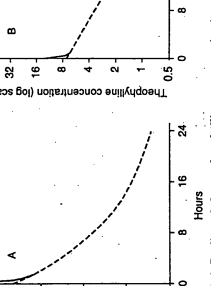
centration-time profiles in the first 2 h is explained by the time required to distribute The discrepancy between the observed (solid line) and predicted (broken line) conthe drug throughout the body (distribution

ing aminophylline, must be given by shortterm infusion over ≥ 5 to $1\overline{0}$ min to avoid time, single IV doses of many drugs, includside effects Extravascular: After a single 300-mg oral is not complete at this time.

Constant-Rate Infusion

after an IV infusion of aminophylline at a constant rate of 45 mg/h (see curve A in Fig. 299-3), the plasma concentration and amount of theophylline in the body increase rate. The plasma concentration and the clearance and elimination rate constant (see Table 299-1), infusion rate equals clearance times plateau plasma drug concentration or until the elimination rate equals the infusion amount of drug in the body are then at steady state—a plateau. Based on the formulas for equals elimination rate constant times pla-





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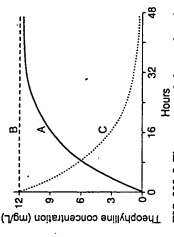
Theophylline concentration (mg/L)

FIG. 299-1. Decline of plasma theophylline concentration in patient A after IV admin**istration of a single 320-mg dose of aminophylline.** Shown on linear (A) and semilogarithmic (B) plots. (blactved curve = (----); predicted curve from given parameter values = (----).

Theophylline concentration (mg/L) phase). Because drug distribution requires

to patient A (see Fig. 299-2), the time course The more rapid the absorption, the closer the ten used orally, is 85% theophylline) is given 209-1) because time is required to absorb the drug. However, AUC is the same because curve is to that of the IV dose. The time of peak concentration is when the absorption dose of aminophylline (anhydrous form, ofrate equals the elimination rate; absorption differs from that of a single IV dose (see Fig. this drug is virtually completely absorbed

Plateau concentration: In patient



ä

stant-rate IV infusion of aminophylline without and with a 530-mg IV loading dose in patient A. A = without loading dose; B = with loading dose; C = drug remaining from ylline concentration after a 45-mg/h con-FIG. 299-3. Time course of plasma theoph loading dose

FIG. 299-2. Time course of plasma theophtion of a single 300-mg dose of aminophyl-

Hours

ylline concentration after oral administra-

line to patient A.

tively, as for IV infusions. Bioavailability is an additional factor applicable to extravasit depend on clearance and half-life, respeccular administration.

only by clearance and infusion rate, and the

teau amount of drug in the body. Thus, the plateau plasma concentration is determined plateau amount of drug in the body is deter-

mined only by the elimination rate constant

and infusion rate.

quired for theophylline to accumulate in the body (and then to disappear) depends on the drug's half-life, as shown in Fig. 299-3. A single IV 530-mg bolus of aminophylline pro-

Time to reach plateau: The time re-

Multiple Oral Doses

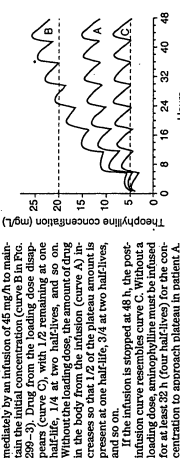
tration of aminophylline 300 mg po q 6 h to tration (see curve A in Fig. 299-4). As with Drug accumulation: Repetitive adminispatient A increases the theophylline concen-

duces a theophylline concentration of 12

mg/L (67 μ mol/L); the bolus is followed immediately by an infusion of 45 mg/h to maintain the initial concentration (curve B in Fig.

299–3). Drug from the loading dose disappears (curve C), with 1/2 remaining at one Without the loading dose, the amount of drug

half-life, 1/4 at two half-lives, and so on. in the body from the infusion (curve A) in-



after oral administration of aminophylline 300 mg q 6 h. Curve A = patient A; curve B FIG. 299-4. Accumulation of theophylline " patient B, whose clearance is 1/2 that of since is twice that of patient A. The dashed lines are the usual therapeutic limits, repreputient A; rurve C = patient C, whose clearnenting the therapeutic window.

after the plateau estimates theophylline

The principles for IV infusion apply to any constant-rate input (eg, to constant-rate de-

clearance.

A plasma concentration measurement made

infusion curve resembles curve C. Without a

for at least 32 h (four half-lives) for the concentration to approach plateau in patient A

If the infusion is stopped at 48 h, the post-

and so on.

and intrauterine drug delivery). Plateau

vices used in transdermal, intraocular, oral,

plusma concentration and the time to reach

IV infusion, the average plateau concentration depends on clearance, and the time required for the drug to accumulate depends on half-life. Here, however, plasma concentrations fluctuate because dosing is intermit-

and C). Patient B has heart failure with a If theophylline clearance is altered, eg, by disease, pharmacokinetics change (curves B because half-life (16 h) is twice that in a clearance of 21.5 mL/h/kg (about half that of patient A). After patient B is given aminophylline 300 mg q 6 h, drug concentration is double that of patient A (curve B), and the time to reach plateau levels is twice as long healthy adult. Plasma theophylline concentrations of 10 to 20 mg/L (55 to 110 µmol/L) is more likely. Thus, patient B is at risk of heart failure decreases metabolism, may be averted by giving a smaller dose. Also, slow metabolism may be detected by monitoring are usually optimal. Above 20 mg/L, toxicity toxicity (nausea, vomiting, CNS stimulation, seizures), which, with the knowledge that plasma concentration.

phylline 200 mg q 8 h (25 mg/h) is probably appropriate. However, because of the long idly produce a therapeutic concentration (and response). The required loading dose of aminophylline is the apparent volume of distribution times the desired theophylline con-Dosage regimens: For patient B, aminohalf-life and the slow accumulation in this patient, a loading dose must be given to rapcentration, corrected by the fraction of the ophylline in aminophylline, or about 500 mg:

 $35 L \times \frac{12 \text{ mg}}{100 \text{ mg aminophylline}} \times \frac{100 \text{ mg aminophylline}}{100 \text{ mg aminophylline}}$ 85 mg theophylline

sustained, 600 mg q 6 h will probably prevent. In a young, otherwise healthy asthmatic theophylline clearance is 86 mL/h/kg, and half-life is 4 h. Aminophylline 300 mg q 6 h (50 mg/h) is probably ineffective (see curve C in Fig. 299-4). The need for more drug can measuring plasma concentration just before the next dose. However, giving aminophylline to this patient is difficult because of the short half-life, high clearance, and large dosdicated. Because absorption is more or less be anticipated and may be confirmed by tient, a prolonged-release formulation is inadult who is a heavy smoker (patient C), age requirements (100 mg/h). For this paconcentrations from fluctuating widely.

PARAMETER VALUES **VARIABILITY IN**

Many factors affecting pharmacokinetic parameters should be considered when tailoring drug administration for a particular patient. Even with dosage adjustment, however, sufficient variability usually remains; thus, drug response and, in some cases, plasma drug concentration must be closely monitored.

Age and weight: For some drugs, the effects of age and weight on pharmacokinetics to 20 yr, renal function appears to correlate are well established. For persons aged 6 mo well with BSA. Thus, for drugs primarily BSA also correlates with metabolic clearcommon. For newborns and infants, renal oped, and generalizations, except for the ocrenal function decreases about 1%/yr. Thus, dosage of these drugs can be adjusted by age. ance in children, although exceptions are and hepatic functions are not fully develeliminated unchanged by renal excretion clearance in children varies with age according to change in BSA. For persons > 20 yr currence of rapid change, are less accurate.

clearance of most drugs appears to vary directly with creatinine clearance, regardless Renal function impairment: Renal of which renal disease is present. The change in total clearance depends on the contribution of the kidneys to total elimination. Thus, nal function (creatinine clearance) for drugs total clearance should be proportional to reexcreted unchanged and to be unaffected for drugs eliminated by metabolism.

Renal failure may change the apparent volume of distribution, which decreases for digoxin because of decreased tissue binding and many other drugs because of decreased and increases for phenytoin, salicylic acid binding to plasma proteins.

Consequently, the binding of several drugs Physiologic stress: Concentration of the acute-phase protein α₁-acid glycoprotein increases during physiologic stress (eg, MI, surgery, ulcerative colitis, Crohn's disease). (eg, propranolol, quinidine, disopyramide) volume of distribution of these drugs deto this protein increases, and the apparent creases accordingly.

Hepatic disease: Hepatic dysfunction can change metabolic clearance, but good correlates or predictors of the changes are

unavailable. Hepatic cirrhosis can dramatically reduce drug metabolism and often results in reduced plasma protein binding because of lowered plasma albumin. Acute hepatitis, with elevated serum enzymes, usually does not after drug metabolism.

Other diseases: Heart failure, pneumonia, hyperthyroidism, and many other diseases can alter the pharmacokinetics of

may be affected by drug interactions. Most interactions are graded, and the extent of the both drugs. Thus, determining and adjusting Drug interactions: Pharmacokinetic parameter values and, therefore, drug response interaction depends on the concentrations of drug dosage is difficult (see Drug Interacnons in Ch. 301).

Dosage: In some instances, changes in dose, dosing rate, or duration of therapy alter

a drug's kinetics. For example, as dose is decreases because of the drug's low solubility in the fluids of the upper GI tract. For increased, the bioavailability of griseofulvin phenytoin, steady-state plasma concentration increases disproportionately when dosing rate is increased, because the metabolizing enzyme has a limited capacity to eliminate the drug, and the usual dosing rate lism. Plasma carbamazepine concentration Other causes of dosage-dependent kinetic tissue binding (eg, phenylbutazone), saturable secretion in the kidneys (eg, high-dose penicillin), and saturable metabolism during approaches the maximum rate of metabodecreases during long-term use because carbamazepine induces its own metabolism. changes are saturable plasma protein and the first pass through the liver (eg, propran-

/ PHARMACODYNAMICS 300 /

Study of the biochemical and physiologic effects of drugs and their mechanisms of action. teins, on or within cells. Some drug classes Many drugs produce pharmacologic reinteract with nucleic acids, metal chelating sponses by interacting with (binding to) specific macromolecules, usually complex proreact directly with endogenous or exogenous nonprotein substances; included are some cancer chemotherapeutic drugs that drugs (eg, calcium disodium edetate, dimer-

DRUG-RECEPTOR INTERACTIONS

caprol, deferoxamine), and antacids used to

chemically neutralize gastric acid.

pine inhibits the actions of acetylcholine on exocrine glands and smooth muscles, but not (hormone, neurotransmitter, intracellular Few if any drugs have absolute specificity, but most have relative selectivity; eg, atrocal binding to cellular components called receptors. Physiologic receptors are macromolecules involved in chemical signaling between and within cells. A molecule that binds on skeletal muscle. The action of such selective drugs results from their physicochemito a receptor is called a ligand. When a ligand

through guanine nucleotide-binding proteins (G proteins) to various effector systems combines with a receptor, cell function changes (see TABLE 300-1). Each ligand may interact with multiple receptur subtypes. Activated receptors directly or indirectly regulate cellular biochemical processes (eg, ion conductance, protein phosphorylation, DNA transcription). In many cases, receptors within the cell membrane are coupled involving intracellular second messenger messenger molecule, or exogenous drug) molecules.

Receptors are dynamic, influenced by exulatory mechanisms. Receptor up-regulation clinically important adaptation to drugs (desensitization, tachyphylaxis, tolerance, acternal factors as well as by intracellular regquired resistance, postwithdrawal supersendown-regulation are relevant sitivity).

lar regions of receptor macromolecules to which Ilgands bind. A drug may interact at the sume site as an endogenous agonist (hor-Recognition sites are the precise molecumone or neurotransmitter) or at a different site. Agonists that bind to an adjacent or a different site are sometimes termed allo-

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